App Serial # 09/714,883 Turner & Mathur Exhibit B LEX-0092-USA

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Novel Human Secreted Proteins and Polynucleotides Encoding the Same
    SEP 2 & 2002
FASTA searches a protein or DNA sequence data bank
 version 31.3t05 March 30, 2000
Please cite:
W.R. Pearson & D.J. Lipman PNAS (1988) 85:2444-2448
/tmp/fastaCAAQPaiHi: 995 aa
 >SEQ ID NO:2
 vs /tmp/fastaDAARPaiHi library
searching /tmp/fastaDAARPaiHi library
                      1 sequences
   1008 residues in
FASTA (3.34 January 2000) function [optimized, BL50 matrix (15:-5)] ktup: 2
 join: 39, opt: 27, gap-pen: -12/ -2, width: 16
 Scan time: 0.050
                                                        opt
The best scores are:
NM_000096 ACCESSION:NM_000096 NID: gi 4557484 ref (1008) 2671
>>NM_000096 ACCESSION:NM_000096 NID: gi 4557484 ref NM_0 (1008 aa)
 initn: 1414 init1: 972 opt: 2671
Smith-Waterman score: 3980; 57.002% identity in 1014 aa overlap (1-992:1-1008)
                                                    50
                       20
                                 30
              10
      MKILILGIFLFLCSSPGWAIDRHCYIGIEESIWNYAPSGKNMLNEKPFSEDLE----FLQ
SEO
      .. ..: .: :
NM_000 MKILILGIFLFLCSTPAWAKEKHYYIGIIETTWDYA---SDHGEKKLISVDTEHSNIYLQ
                                             40
                                 30
              10
                       20
                                     90
                                             100
        60
                  70
                           80
      GGQARKSFVFKKALYFQYTDNTFQRIIEKPSWLGFLGPMIKAETGDFIYVHVKNNASRAY
SEO
       NM_000 NGPDRIGRLYKKALYLQYTDETFRTTIEKPVWLGFLGPIIKAETGDKVYVHLKNLASRPY
                                    90
                                            100
                                                      110
       60
                 70
                          80
                                    150
                                             160
                130
                          140
       120
      SYHPHGLTYSKENEAHGAIYPDNTTGLQKEVEYLEPGKQYTYKWYVEEHQGPGPNDSNCV
SEO
       ... ::.:: ::.: ::::::::: .:. . . . ::.:::: . . :.::::
NM_000 TFHSHGITYYKEHE--GAIYPDNTTDFQRADDKVYPGEQYTYMLLATEEQSPGEGDGNCV
                                              160
                                     150
      120
                130
                           140
                                    210
                                             220
                          200
                190
       180
      TRIYHSHIDTARDVASGLIGPILTCKRGTLNGDTEKDIDRSSFLMFSTTDESRSWYSDEN
SEO
      .:::..:: ::: ..:
NM_000 TRIYHSHIDAPKDIASGLIGPLIICKKDSLDKEKEKHIDREFVVMFSVVDENFSWYLEDN
                                              220
                 190
                           200
                                     210
        180
                           260
                                     270
                                              280
                                                        290
       240
                 250
```

IRAF-TESGKINTSDPRFEESMSMQSINGYIYGNLPNLTMCAEDRVQWYFVGMGGVADIH

PVYLRGQTLISRNHRKDTIMLFPSSLEDAFMVAKAPGVWMLGCQ----IHESMQAFFKVS

270

330

330

280

340

340

350

NM\_000 IKTYCSEPEKVDKDNEDFQESNRMYSVNGYTFGSLPGLSMCAEDRVKWYLFGMGNEVDVH

NM\_000 AAFFHGQALTNKNYRIDTINLFPATLFDAYMVAQNPGEWMLSCQNLNHLKAGLQAFFQVQ

260

320

320

SEO

SEO

240

300

300

250

310

310

SEQ	360		380	390	400 KKNLTAAGSKS	410 SQLFFERSPTR
~	. : . :				:: :	.::: ::
NM_000	ECNKSSSKDN: 360	RGKHVRHYYI 370			KENLTAPGSDS 400	SAVFFEQGTTR 410
	420	430	4	40 4	150 4	160
SEQ	IGGTYKKLIY	REYTDASFQTQ	KAREEH	LGILGPVFKA	AEVGQTIKITE	FYNNASLPLSI
NM_000	:::.:::::: IGGSYKKLVYI					CHNKCANDIGT
NM_000	420		440	450	460	470
4	470 48			500		520
SEQ	QPPGLHYNKS					
NM_000	EPIGVRFNKNI 480	NEGTYYSPNYN		ASHVAPTETE		
SEQ	530 TWFYYSSVNG			560 GDDGKQKGVI		580 DENESNLLDE
	:::.:.					
NM_000	AKMYYSAVDP1 540	SKDIFTGLIGF 550	MKICKKGSL 560	HANGRQKDVL 570		590
SEQ	590 N-RTFITEPEN					
NIM OOO	: : : :	.:::: : :	SNKMHSMNG	.:::: ::: FMYGNOPGLT	: :::::: MCKGDSVVWY	.::.:. :. LFSAGNEADV
1411_000	600	610			640	650
	650				690	
SEQ	HGIYFSGNTFT					LGGMKHKYHV
NM_000	:::::::: HGIYFSGNTYI 660	WRGERRDTAN		HMWPDTEGTF		
SEQ	710 RQCGKPNPDQT	QYQEEKIIIT		YSPSRKWENE	LHHLRRENQT	
NIM OOO	:: : .: NQCRRQSEDST	: :. FVI.GERTVV-				
1111_000	720	730	740	750	760	770
SEO	770 LGSKYKKVLYF	780	790	800	810	820 KNKAARPYST
SEQ	.:::::::::				. ::::	
NM_000	IGSKYKKVVYR 780		VERKAEEEHI 800	LGILGPQLHA 810	DVGDKVKIIF 820	KNMATRPYSI 830
	830	840	850	860	870	880
SEQ	HAHGVKTNNST	VVPTQPGEIQ				
NM_000	::::::::::::::::::::::::::::::::::::::		:.:.:: TYVWKIPER: 860			.:::.::. QVKDLYSGLI 890
	5 10			= · <del>*</del>	<del>-</del>	•

GPLSVCR----KDINPN-IVHRVLHFMIFDENESWYFEDSINTYASKPNKVDKENDNFQL SEQ NM\_000 GPLIVCRRPYLKVFNPRRKLEFALLFLVFDENESWYLDDNIKTYSDHPEKVNKDDEEFIE SNQMHAINGRLFGNNQGITFHVGDVVNWYLIGIGNEADLHTVHFHGHSFEYKHKYLI SEO NM\_000 SNKMHAINGRMFGNLQGLTMHVGDEVNWYLMGMGNEIDLHTVHFHGHSFQYKHR 

995 residues in 1 query sequences 1008 residues in 1 library sequences Scomplib [version 3.3t05 March 30, 2000] start: Wed Sep 18 11:16:31 2002 done: Wed Sep 18 11:16:32 2002 Scan time: 0.050 Display time: 1.417

Function used was FASTA